

## COMMUNICATION

## Zinc-Catalyzed Synthesis of 2-Alkenylfurans via Cross-Coupling of Enynones and Diazo Compounds

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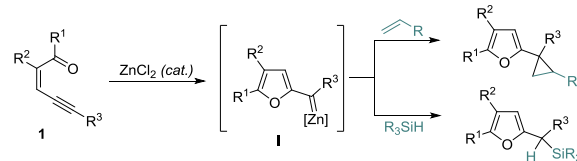
**Inexpensive, low-toxic  $\text{ZnCl}_2$  serves as the catalyst for selective cross-coupling of enynones and diazocompounds, an unprecedented reaction pathway for zinc carbenoids. A cascade sequence comprising formal cyclization/cross-coupling leads to a variety of 2-alkenylfurans.**

The replacement of precious metals with catalysts based on more abundant and low-toxic metals is gaining relevance due to economic and environmental reasons, both in industry and academia.<sup>1</sup> As a result, elegant transformations based on the use of iron, copper or cobalt catalysts, among others, have appeared in the recent years.<sup>2</sup> Whilst the utility of zinc in synthetic organic chemistry is known for more than a century, the development of zinc-catalyzed transformations remains less developed.<sup>3,4</sup> For instance, the well-known Simmons-Smith cyclopropanation is typically accomplished with over-stoichiometric amounts of zinc (moisture and oxygen sensitive  $\text{Et}_2\text{Zn}$ ).<sup>5</sup> With the aim to overcome this limitation, we became interested in exploring the feasibility to generate zinc carbenoids in a catalytic manner and to exploit their reactivities for the synthesis of relevant compounds. Thus, we recently reported the use of enynones **1** as carbene precursors using simple  $\text{ZnCl}_2$  as the catalyst (Scheme 1, a).<sup>6,7</sup> The cyclization of **1** generates a 2-furyl zinc carbenoid intermediate **I**,<sup>8</sup> which can be further trapped with alkenes or silanes to yield valuable 2-cyclopropyl- or 2-alkylsilylfurans, respectively. Computational studies provided support for the proposed 2-furyl zinc Fischer-type carbenoid intermediate.<sup>6</sup> This approach provides a catalytic alternative to previously established zinc-mediated cyclopropanation<sup>5b</sup> or Si–H insertions.<sup>9</sup>

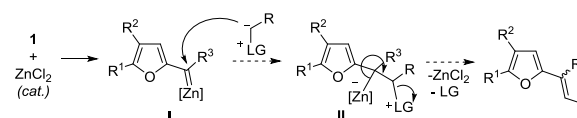
In order to further explore the synthetic potential of the postulated 2-furyl zinc-carbenoid intermediate **I**, we wondered whether the reactivity could be expanded beyond its classical known reactivity patterns. According to the predicted electrophilicity of the putative zinc carbenoid, we envisioned a route to relevant 2-alkenylfuran derivatives<sup>10</sup> by trapping the carbenoid with a suitable carbon-nucleophilic reagent bearing a leaving group. Thus, the formation of

a new C–C bond by nucleophilic attack should set the stage for an elimination process delivering the desired target and allowing catalyst turnover (Scheme 1, b). Herein, we disclosed our findings regarding this unprecedented reactivity of zinc carbenoids.

(a) Zinc-carbenoid catalytic generation: Classical reactivity (Ref. 6a)



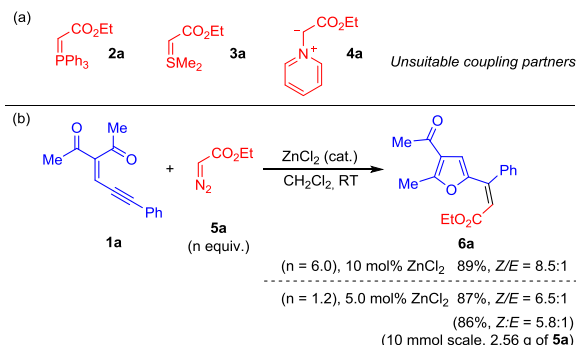
(b) Zinc-carbenoid catalytic generation: Unprecedented reactivity (this work)



**Scheme 1.** Reactivity of in situ generated 2-furyl zinc carbenoids.

The evaluation of the initial hypothesis was carried out with readily available enynone **1a** as benchmark reagent and  $\text{ZnCl}_2$  as the catalyst. As coupling reagent, we selected stabilized ylides **2-4a** (Scheme 2, a) which meet the requirements to achieve the desired transformation since they are good nucleophiles and contain a leaving group. Unfortunately, only phosphorous ylide **2a** led to the formation of the desired 2-alkenylfuran, albeit in unpractical low yield (<10%). Then, we turned our attention to stabilized diazo compounds. In contrast to other transition metals, in general, stabilized diazo compounds do not decompose toward carbenoids in the presence of zinc salts,<sup>11</sup> yet they are good carbon nucleophiles and have an excellent leaving group as  $\text{N}_2$ .<sup>12</sup> Indeed, using ethyl diazoacetate (EDA, **5a**) under reaction conditions similar to those previously reported for related transformations (10.0 mol%  $\text{ZnCl}_2$ , 6.0 equiv. of **5a**, 25 °C,  $\text{CH}_2\text{Cl}_2$ , 1 h),<sup>6a</sup> we isolated 2-alkenylfuran **6a** in good yield and selectivity (89%, Z:E = 8.5:1) (Scheme 2, b). A

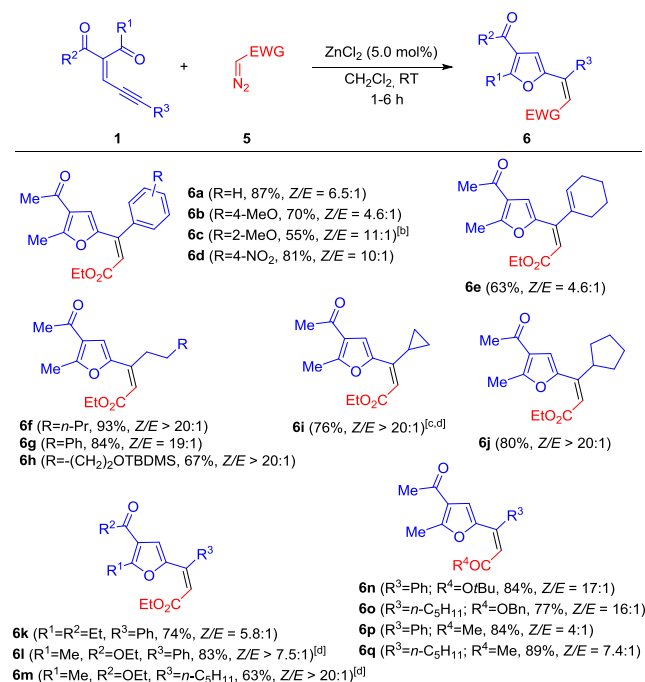
further optimization demonstrated that 5.0 mol% of  $\text{ZnCl}_2$  and only a slight excess of **5a** (1.2 equiv.,  $\text{CH}_2\text{Cl}_2$ , 25 °C, 1 h) gave rise to comparable results (87%,  $Z:E = 6.5:1$ ).<sup>13</sup> Additionally, to check the practicality of this protocol, the test reaction was carried out at larger scale (10 mmol) with almost the same efficiency (2.56 g of **6a**, 86%,  $Z:E = 5.8:1$ ). The formation of compound **6a** validates our initial hypothesis as it represents an *unprecedented reaction pathway for zinc carbenoids*. Besides, the overall transformation comprises a rather unusual example of formal selective hetero-coupling of two different carbenoid precursors.<sup>14</sup> Notably,  $\text{ZnCl}_2$  showed better efficiency and selectivity when comparing it to other representative metal catalysts known capable of generating carbenoids from enynes.<sup>15</sup>



**Scheme 2.** Zinc-catalyzed preparation of 2-alkenylfuran **6a**: Initial findings and optimized reaction conditions.

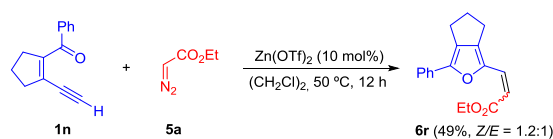
With the optimized reaction conditions in hand, we next explored the scope of this transformation (Table 1). First, the reaction of EDA (**4a**) with an array of easily available enynones **1** was examined. Thus, reactions with aryl substituted enynones proceeded smoothly to afford compounds **6a-d**, and those with both electron-rich and electron-deficient aryl groups gave the expected products in good yields and variable stereoselectivity. Even an *ortho*-substituent in the aryl group is well tolerated, though a slight decrease of the yield was observed.<sup>16</sup> An alkenyl substituent at the alkyne terminus was also suitable giving rise to compound **6e** (63%,  $Z:E = 4.6:1$ ). Interestingly, alkyl substituents on the alkyne, both primary and secondary, afforded efficiently the corresponding 2-alkenylfurans **3f-6j** with high  $Z$ -selectivity.<sup>17</sup> Notably, the presence of a protected alcohol in the substrate was compatible with this protocol affording the furan derivative **6h** in a reasonable yield (67%) and a high  $Z$ -selectivity. Further modifications on the substituents  $R^1$  and  $R^2$  in the enynone led to compounds **6k-m**. Then, we evaluated the diazo compound component. Various carboxylic esters as well as ketones as stabilizing group of the diazo compound proved suitable substituents, yielding furans **6n-q** in respectable yields (77-89%). On the contrary, this procedure could not be applied to phenyl-, trimethylsilyl diazomethane or ethyl 2-diazo-2-phenylacetate, which gave rise to complex reaction mixtures, under various similar reaction conditions tested, while dimethyl 2-diazomalonate proved unreactive.

**Table 1.** Zinc-catalyzed synthesis of 2-alkenylfurans **6**: Scope.<sup>[a]</sup>



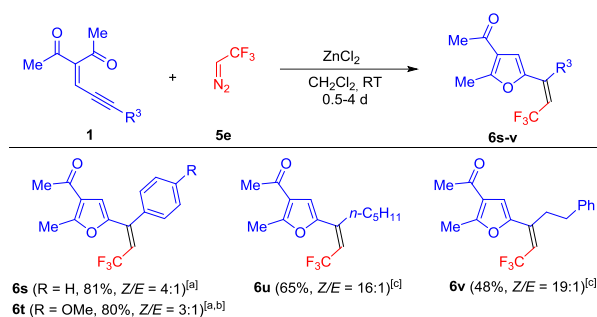
<sup>[a]</sup> Reaction conditions: **1** (0.2-0.3 mmol), **5** (1.2 equiv.),  $\text{ZnCl}_2$  (5.0 mol%),  $\text{CH}_2\text{Cl}_2$  (0.1 M), RT. Yields are referred to isolated products.  $Z:E$  Selectivity was determined by NMR. [b] See Ref. 16. [c] At 0 °C. [d] <5% of an unidentified isomer was detected.

Interestingly, we found that other enynones such as compound **1n** proved applicable to this coupling. Indeed, treatment of **1n** with EDA (**5a**) in the presence of 10 mol% of  $\text{Zn}(\text{OTf})_2$  resulted in the formation of tetrasubstituted furan **6r** in moderate yield (49%) after stirring in 1,2-dichloroethane at 50 °C for 12 h (Scheme 3).



**Scheme 3.** Zinc-catalyzed preparation of 2-alkenylfuran **6r** from enynone **1n**.

The preparation of chemicals bearing a trifluoromethyl group has recently attracted great interest in synthesis. Since trifluoromethyl diazomethane can be conveniently generated in situ,<sup>18</sup> we checked the feasibility to access the corresponding alkenyl- $\text{CF}_3$  furyl derivatives as a complementary approach towards this relevant moiety.<sup>19</sup> Indeed, the reaction of various representative enynones **1** with an excess of trifluoromethyl diazomethane (**5e**, ~20 equiv.) generated from trifluoromethylamine hydrochloride and  $\text{NaNO}_2$ , in the presence of  $\text{ZnCl}_2$  allowed us to obtain  $\text{CF}_3$ -substituted alkenes **6s-v** in synthetically useful yields (Scheme 4).<sup>20</sup>



**Scheme 4.** Zinc-catalyzed synthesis of CF<sub>3</sub>-substituted 2-alkenylfurans **6s-v**. Yields are referred to isolated product. Z/E selectivity was determined by NMR. <sup>[b]</sup> ZnCl<sub>2</sub> 100 mol%. <sup>[b]</sup> At 0 °C. <sup>[c]</sup> ZnCl<sub>2</sub> 20 mol%.

As depicted in Scheme 1 (b), the formation of 2-alkenylfurans **6** can be explained through the initial 5-exo-dig cyclization of enynones **1** to generate the 2-furyl zinc carbenoid species **I**. Subsequent nucleophilic attack of the diazocompound **5** likely afford intermediate **II**.<sup>12</sup> Favoured extrusion of N<sub>2</sub> with a concomitant release of the zinc catalyst leads to furans **6** and allows catalyst turnover.<sup>21,22</sup>

## Conclusions

In summary, we have herein reported a new zinc-catalyzed formal selective hetero-coupling of two carbenoid precursors as enynones **1** and diazocompounds **5**. The observed reactivity constitutes an unprecedented reactivity pattern for zinc carbenoids. This transformation has enabled the preparation of valuable 2-alkenylfuran derivatives **5** with a remarkable scope.<sup>10</sup> The overall process comprises a cyclization of the enynone **1** followed by a coupling with the diazocompound and means the formation of new C–O and C=C bonds. Relevant CF<sub>3</sub>-substituted 2-vinylfuran analogues can also be accessed with this methodology using in situ generated trifluoromethyl diazomethane. The use of an inexpensive and low-toxic ZnCl<sub>2</sub> as catalyst is remarkable within the context of the new venues of chemistry regarding the development of sustainable transformations.

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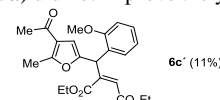
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† Electronic Supplementary Information (ESI) available: Experimental procedures, characterization data and NMR spectra. See DOI: 10.1039/c000000x/

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- Experiments were carried out using CuBr and [(IPr)Au(NTf<sub>2</sub>)] under otherwise identical reaction conditions: CuBr (**6a**, 43%, Z/E = 1.4:1, CH<sub>2</sub>Cl<sub>2</sub>, 25 °C, 24 h); [(IPr)Au(NTf<sub>2</sub>)] (**6a**, 71%, Z/E = 1.7:1, CH<sub>2</sub>Cl<sub>2</sub>, 25 °C, 24 h).
- Enynone **1c** bearing *o*-MeO-C<sub>6</sub>H<sub>4</sub> group led to the formation of compound **6c'** (11%) as byproduct. The use of a larger excess of ethyl diazoacetate (**5a**) did not improve the yield of **6c'**.



- An enynone bearing a bulky R<sup>3</sup> group such as *tert*-butyl proved unreactive towards **2a** under several reaction conditions.
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- 20 See ESI for specific conditions on the generation of **2e** and the reaction conditions for the formation of **3s-v**.
- 21 Preliminary computational studies indicate that (*E*)-**3** are the most stable isomers. Thus, without further additional evidences, we believe that *Z/E* selectivities likely originate from kinetic control. See the ESI for computational details.
- 22 At this stage, the following pathway cannot be ruled out: *i*) activation of the substrate by coordination of the zinc to the carbonyl group(s); *ii*) conjugate attack of the diazo compound to the triple bond; *iii*) cyclization with elimination of N<sub>2</sub>. However, other common Lewis acids tested (BF<sub>3</sub>·Et<sub>2</sub>O, AlCl<sub>3</sub>, TiCl<sub>4</sub>, MgCl<sub>2</sub>, under catalytic or stoichiometric conditions) did not give rise to the expected 2-alkenylfuran derivative **5**. In contrast, other metals known to activate alkynes to generate carbenoids as Cu or Au showed the same reactivity, see Ref. 15.